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(54) CEREBRAL ORGANIC ANION TRANSPORTER AND ITS GENE

(57) A cerebral organic anion transporter OAT3 which is useful as a protein regulating the uptake/excretion of organic anionic substances in the brain; a nucleic

acid having a base sequence encoding the same; and an antibody against the same. The amino acid sequence and the base sequence of the above OAT3 are shown in Sequence Listing in the description.

EP 1 114 830 A

Description

Technical Field

[0001] The present invention relates to a gene involved in organic negative ion (organic anion) transport and the polypeptide encoded by the gene.

Background Art

- [0002] Liver and kidney play important roles in the metabolism and excretion of biologically foreign compounds and drugs out of bodies. Tubule cells and hepatocytes belong to epithelial cells with polarities. It is supposed that some of anionic substances are taken up through the basolateral membranes into kidney and liver by transporters, while the organic anions generated metabolically in cells are excreted by transporters.
 - [0003] The uptake of organic anions through the basolateral membranes of tubule cells and hepatocytes have been investigated so far in experiment systems using isolated organ perfusion protocols, dissected cells and membrane vesicles. According to such conventional methods, however, the detailed analysis of the transport of organic anions through the basolateral membranes has been difficult. Accordingly, it has been desired to isolate the transporters per se and analyze the properties of transporters in detail.
 - [0004] Alternatively, plural experimental results suggestively indicate the presence of the transport of organic anions in brain. The transport of organic anions in brain is supposed to function for the extracerebral excretion of endogenous and exogenous organic anions.
 - [0005] Although the transport of organic anions in brain is speculated to play an important role in the elimination of endogenous anions and foreign compounds from brain, the detail of the transport therein is more ambiguous than the transport in kidney and liver, due to the difficulty in physiological experiments therein.
 - [0006] Based on these backgrounds, the screening of the organic anion transporter molecules per se has been actively carried out in 1990 and thereafter. Consequently, two organic anion transporters derived from the basolateral membrane of liver have been isolated until the last year. (Hagenbuch, B. et al., Proc. Natl. Acad. Sci. USA, Vol. 88, pp. 10629-33, 1991; Jacquemin, E. et al., Proc. Natl. Acad. Sci. USA, Vol. 91, pp. 133-7, 1994)
 - [0007] The present inventors independently isolated an organic anion transporter OAT1 responsible for the most important function in the organic anion transport in kidney successfully last year (Sekine, T., et al., J. Biol. Chem., Vol. 272, pp. 18526-9, 1997) and already filed the patent application thereof. OAT1 is a transporter capable of transporting a great number of organic anions with different chemical structures and is also involved in the transport of various anionic drugs. OAT1 is expressed in a specific manner to kidney, while OAT1 is very slightly expressed in brain except kidney.
 - [0008] Recently, the inventors have further identified a liver-specific organic anion transporter (OAT2) with about 40 % homology to OAT1 in terms of amino acid level (FEBS letter, Vol. 429, pp. 179-182, 1998) (Japanese Patent Application No. 169174/1998).
 - [0009] The isolation and identification of OAT1 and OAT2 indicates that these organic anion transporters form a family. Additionally because OAT2 is expressed specifically in liver, it is suggested that the family is not kidney-specific but is expressed in various organs.
 - [0010] As described insofar, it is suggested that an organic anion transport system is present in brain, but the OAT1 expression in brain is quite slight while OAT2 is not present therein. Based on these findings, the Inventors have anticipated the presence of an unknown transporter responsible for the organic anion transport in brain.
 - [0011] Alternatively, the organic anion transport in the basolateral membrane of liver is complicated; particularly, the efflux flow of conjugated substances (many of the conjugated substances are organic anions) generated at a vast scale in hepatocytes into blood has not yet been known. The organic anion transport in liver cannot sufficiently be described on the single basis of the organic anion transporters including OAT2. Hence, the presence of an unknown transporter is suggested.
 - [0012] The inventors isolated the organic anion transporter OAT1 serving as the most important role in the organic anion transport in kidney (Sekine, T. et al., J. Biol. Chem. Vol. 272, pp. 18526-9, 1997). Based on the structural similarity to OAT1, the inventors identified a liver-specific organic anion transporter (OAT2) (Sekine, T., et al., FEBS letter, Vol. 429, pp. 179-182, 1998). The inventors already reported additionally (Sekine, T., et al., J. Biol. Chem., Vol. 272, pp. 18526-9, 1997) that OAT1 had low homology to an organic cation transporter OCT1 (Grundemann, D. et al., Nature, Vol. 372, pp. 549-52, 1994).
 - [0013] Taking account of these evidence, the inventors Identified a sequence common to OAT1, OAT2 and OCT1 and prepared a degenerate primer based on the sequence. By using the degenerate primer, the inventors identified a novel cDNA fragment with low homology to OAT1, OAT2 and OCT1 from rat brain mRNA by RT (reverse transcript)-PCR (polymerase chain reaction) method. By using the cDNA fragment, a cDNA never reported yet was discriminated

from the rat cDNA library. The resulting protein was designated cerebral type organic anion transporter OAT3 as a third member of the OAT family.

Disclosure of the Invention

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[0014] The invention relates to the organic anion transporter OAT3. The inventive organic anion transporter OAT3 is a transporter with a wide range of substrate selectivity and transports organic anions with different chemical structures (having a potency to take up the organic anions). However, no substantial uptake of a typical organic cation TEA (tetraethylammonium) is observed. Hence, the inventive organic anion transporter OAT3 with a wide range of substrate selectivity is an organic anion transporter with no substantial substrate selectivity of TEA (tetraethylammonium) as the typical organic cation but is selectively distributed in organs mainly including brain and liver.

[0015] The inventive protein includes the organic anion transporter OAT3 of an amino acid sequence represented by SQ ID No. 2 (in human) or 4 (in rat) or of an amino acid sequence with such a modification of the aforementioned amino acid sequence as deletion, substitution or addition of one or several amino acids. The deletion, substitution or addition of amino acids is satisfactory at an extent such that no organic anion transport activity is deteriorated; the number of the amino acids then is generally 1 to about 110, preferably 1 to about 55. Such protein has generally 60 to 80 %, preferably 70 to 90 % homology in amino acid sequence to the amino acid sequence represented by SQ ID No. 2 or 4.

[0016] Furthermore, the invention encompasses a nucleic acid, preferably DNA or RNA, encoding the inventive protein comprising the organic anion transporter OAT3. The inventive nucleic acid encompasses the nucleic acid encoding the inventive protein and nucleic acids hybridizable with the nucleic acid under stringent conditions.

[0017] Still furthermore, the invention relates to a partial sequence of the nucleic acid encoding the inventive protein or nucleotides hybridizable with the partial sequence under stringent conditions.

[0018] Still yet furthermore, the invention relates to an antibody against the inventive protein or a polypeptide immunologically identical to the inventive protein.

Brief Description of the Drawings

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- Fig. 1 depicts the organic anion uptake activity of the inventive rat OAT3 expressed in Xenopus oocyte;
- Fig. 2 depicts the results of kinetic analyses of the transport of PAH, estrone sulfate and ochratoxin A with the inventive rat OAT3 in the oocyte;
- Fig. 3 depicts the results on the inhibition of the organic anion transport with the inventive rat OAT3 by various organic substances;
- Fig. 4 depicts the results of the Northern blotting analysis of the inventive rat OAT3 gene;
- Fig. 5 depicts the results on the inhibition of the rat OAT3 transport by various metabolites of cerebral type neurotransmitters;
- Fig. 6 depicts the uptake activity of ¹⁴C-PAH (p-aminohippuric acid) when the inventive hOAT3 was expressed in Xenopus oocyte;
- Fig. 7 depicts the uptake activity of ³H-estrone sulfate when the inventive hOAT3 was expressed in Xenopus oocyte; Fig. 8 depicts the uptake activity of ³H-dehydroepiandrosterone sulfate when the inventive hOAT3 was expressed in Xenopus oocyte;
- Fig. 9 depicts the uptake activity of ³H-ochratoxin A when the inventive hOAT3 was expressed in Xenopus oocyte; Fig. 10 depicts the uptake activity of ³H-cimetidine when the Inventive hOAT3 was expressed in Xenopus oocyte; Fig. 11 depicts the uptake activity of ³H-estradiol glucuronide when the inventive hOAT3 was expressed in Xenopus oocyte;
 - Fig. 12 depicts the uptake activity of ³H-prostaglandin E2 when the inventive hOAT3 was expressed In Xenopus oocyte:
- Fig. 13 depicts the uptake activity of ¹⁴C-taurocholic acid when the inventive hOAT3 was expressed in Xenopus
 - Fig. 14 depicts the uptake activity of ¹⁴C-glutaric acid when the inventive hOAT3 was expressed in Xenopus oocyte;
 - Fig. 15 depicts the uptake activity of ³H-methotrexate when the inventive hOAT3 was expressed in Xenopus occyte;
 - Fig. 16 depicts the uptake activity of ¹⁴C-salicylic acid when the inventive hOAT3 was expressed in Xenopus oocyte;
 - Fig. 17 depicts the uptake activity of ¹⁴C-Indomethacin when the inventive hOAT3 was expressed in Xenopus occyte:
 - Fig. 18 depicts the uptake activity of ¹⁴C-cholic acid when the inventive hOAT3 was expressed in Xenopus oocyte; and

Fig. 19 depicts the results on the inhibition of the transport of ³H-estrone sulfate with the inventive hOAT3 by various organic substances.

Best Mode for Carrying out the Invention

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[0020] The inventive organic anion transporter gene can be isolated and identified by screening of tissues and cells of organs such as kidney and brain in appropriate mammalian animals used as gene sources. The mammalian animals include non-human animals such as dog, cow, horse, goat, sheep, monkey, pig, rabbit, rat and mouse and additionally include human.

[0021] The gene screening and isolation can preferably be carried out by homology screening and PCR screening. The base sequence of the resulting cDNA is determined by a conventional method; the translation region is analyzed; and the amino acid sequence of the protein encoded by the cDNA, namely the amino acid sequence of OAT3, can be determined.

[0022] It is verified for example by the following manners that the resulting cDNA is the cDNA of the organic anion transporter gene, namely that the genetic product encoded by the cDNA is the organic anion transporter. More specifically, the cRNA prepared from the isolated OAT3 gene is integrated and expressed in the occyte; then, the transport (uptake) potency of organic anions in cells is confirmed by assaying the incorporation of an appropriate organic anion as the substrate in cells by the general uptake experiment (Sekine, T., et al., J. Biol. Chem., Vol. 272, pp. 18526-9, 1997). [0023] By applying the same uptake experiment to the expression cell, the transport property and substrate specificity of OAT3 can be examined.

[0024] The SQ ID No. 3 in the sequence listing shows the base sequence of the cDNA of the rat organic anion transporter OAT3 isolated by such method; and SQ ID No. 4 shows the amino acid sequence thereof.

[0025] By using the cDNA of the resulting OAT3 gene for screening an appropriate cDNA library or genomic DNA library prepared by using a different gene source, a homologous gene or chromosomal gene derived from a different tissue or a different biological organism or the homology can be isolated.

[0026] The base sequence of the cDNA of human organic anion transporter OAT3 identified by such method is shown as SQ ID No. 1 and the amino acid sequence thereof is shown as SQ ID No. 2.

[0027] By using a synthetic primer designed on the basis of the base sequence as the base sequence (SQ ID No. 1 or 3) of the inventive gene disclosed or a part of the information thereof, the gene can be isolated from the cDNA library by general PCR.

[0028] DNA libraries such as cDNA library or genomic DNA library or the like can be prepared by the method described in for example "Molecular Cloning; Sambrook, J., Fritsh, E. F. and Maniatis, T. ed., issued by Cold Spring Harbor Laboratory Press in 1989". Otherwise, any existing commercially available library can satisfactorily be used.

[0029] The inventive organic anion transporter (OAT3) can be generated by using for example cDNA encoding the organic anion transporter by genetic recombinant technology. For example, DNA (cDNA and the like) encoding the organic anion transporter is integrated in an appropriate expression vector; and the resulting recombinant DNA can then be transfected in an appropriate host cell. The expression system (host vector system) for polypeptide generation includes for example expression systems of bacteria, yeast, insect cells and mammalian cells. Among them, insect cells and mammalian cells are preferably used for the recovery of the functional protein.

[0030] For the expression of the polypeptide in mammals, for example, the DNA encoding the inventive organic anion transporter is inserted in the downstream of an appropriate promoter (for example, SV40 promoter, LTR promoter, elongation 1α promoter and the like) in an appropriate expression vector (for example, retrovirus vector, papilloma virus vector, vaccinia virus vector, SV40 vector and the like) to construct an expression vector. By subsequently transforming an appropriate animal cell with the resulting expression vector and culturing the transformant in an appropriate culture medium, the objective polypeptide can be generated. The mammalian cell as the host includes monkey COS-7 cell, Chinese hamster CHO cell, human HeLa cell, or cell lines such as kidney tissue-derived primary culture cell, porcine kidney-derived LLC-PK1 cell and opposum kidney-derived OK cell and the like.

[0031] As the cDNA encoding the organic anion transporter OAT3, use can be made of cDNA with the base sequence represented by SQ ID No. 1 or 3; as the cDNA, with no specific limitation to the cDNA described above, additionally, DNA corresponding to the amino acid sequence is designed and used, which can encode the polypeptide. In this case, it is known that each amino acid is encoded by one to 6 types of codons, so codons for use can be selected appropriately. For example, a sequence with higher expression can be designed, in terms of the frequency of codons used by a host for expression. DNA with the designed base sequence can be recovered by chemical DNA synthesis, fragmentation and conjugation of the cDNA, and a partial modification of the base sequence. An artificial partial modification of the base sequence or mutagenesis thereof can be carried out by site specific mutagenesis, by utilizing a primer comprising a synthetic oligonucleotide encoding the desired modification "Mark, D. F., et al., Proc. Natl. Acad. Sci. USA, Vol. 8, pp. 5662-5666, 1984".

[0032] Nucleotides (oligonucl otide or polynucleotide) hybridizable with the inventive organic anion transporter gene

under stringent conditions can be used as probe for detecting the organic anion transporter gene and can also be used for example as antisense oligonucleotide, ribozyme and decoy, so as to modify the expression of the organic anion transporter.

[0033] In accordance with the invention, the term hybridization under stringent conditions generally means hybridization in 5 x SSC or a hybridization solution at a salt concentration equal to the concentration under a temperature condition of 37 to 42 °C for about 12 hours, followed by preliminary rinsing in 5 x SSC or a solution at a salt concentration equal to the concentration and rinsing in 1 x SSC or at a salt concentration equal to the concentration. Higher stringency can be realized by carrying out rinsing in $0.1 \times SSC$ or a solution at a salt concentration equal to the concentration.

[0034] Additionally, the invention relates to a partial sequence of the nucleic acid encoding the inventive protein or nucleotides hybridizable with the sequence under stringent conditions. As such nucleotides, generally, use can satisfactorily be made of nucleotides comprising a partial sequence of consecutive 14 or more nucleotides in series in the base sequence represented by SQ ID No. 1 or 3 or a sequence complementary to the partial sequence; so as to enhance the specificity of the hybridization, a longer sequence, for example a sequence of 20 bases or more or a sequence of 30 bases or more, can satisfactorily be used as such partial sequence. These nucleotides can be labeled, if necessary, with radioactive elements, fluorescent substances or chemiluminescent substances.

[0035] The nucleotides comprising a partial sequence of consecutive 14 or more base in series in the inventive base sequence represented by SQ ID No. 1 or 3 or a sequence complementary to the partial sequence preferably carries the specific base sequence of the base sequence encoding the inventive organic anion transporter OAT3 and can satisfactorily be labeled, if necessary.

[0036] By using the inventive organic anion transporter or a polypeptide immunologically identical thereto, additionally, an antibody can be raised. The antibody can be utilized for detecting or purifying the organic anion transporter. The antibody can be raised, by using the inventive organic anion transporter, a fragment thereof, or a synthetic peptide with a partial sequence thereof or the like as an antigen. The antibody, if polyclonal, can be generated by general methods comprising inoculating such antigen in a host animal (for example, rat and rabbit) and recovering the resulting immunized serum. The antibody, if monoclonal, can be generated by techniques such as general hybridoma method. Further, the inventive antibody is satisfactorily prepared as chimera form or humanized antibody.

Best Mode for Carrying out the Invention

30 [0037] The description is now made in more detail in the following examples, but the examples are in no way of limitation of the invention.

[0038] In the following examples, the Individual procedures followed the methods described in "Molecular Cloning; Sambrook, J., Fritsh, E. F. and Maniatis, T. ed., issued by Cold Spring Harbor Laboratory Press in 1989" or followed the instructions of commercially available kit products if used, unless otherwise stated.

Example 1

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Isolation and analysis of multi-selective organic anion transporter 3 (OAT3) cDNA

[0039] (1) Preparation of degenerate primer based on the base sequence information of OAT1, OAT2 and OCT1 [0040] Based on the base sequence information of OAT1 and OAT2 isolated previously by the inventors and the reported base sequence information of OCT1, degenerate primer was prepared with reference to amino acid sequences in common to these three transporters (amino acids 267-275 and amino acids 447-452 in the amino acid sequence of OAT1).

[0041] From rat brain was extracted total RNA by GITC method; and poly(A) + RNA was then purified by using an oligodT column. From the rat brain poly(A) + RNA was prepared cDNA by using reverse transcriptase; using the resulting cDNA as template, PCR was conducted with the degenerate primer. Consequently, a PCR product of about 550 bp was prepared.

[0042] By using a TA cloning kit (manufactured by Invitrogen Co.), the PCR product was cloned; and some of the base sequence was determined. Consequently, a novel cDNA (B10) with homology at the level of 50 % to OAT1 in terms of amino acid level was recovered.

[0043] A probe prepared by labeling B10 cDNA with ³²P was used for Northern hybridization with poly(A) + RNA extracted from various rat organs. Positive bands were visually detected in the liver, kidney, brain and eyes.

[0044] Because the inventors had an excellent cDNA library of rat kidney, the inventors screened the rat kidney cDNA library by using the B10 probe. Hybridization was promoted overnight in a hybridization solution at 37 °C. Thereafter, the filter membrane was rinsed in 0.1 × SSC/0.1 % SDS at 37 °C. As the hybridization solution, use was made of a buffer, pH 6.5 containing 50 % formamide, 5 × standard saline citrate (SSC), 3 × Denhard solution, 0.2 % SDS, 10 % dextran sulfate, 0.2 mg/ml modified salmon sperm DNA, 2.5 mM sodium pyrophosphate, 25 mM MES, and 0.01

% Antifoam B (manufactured by Sigma, Co.). The clone isolated in λ ZipLox was further subcloned in a plasmid vector pZL by in vivo excision method. Consequently, a novel clone (rk1411) with an organic anion transport activity was recovered (Example 2 below is to be referenced concerning transport function analysis).

[0045] Th base sequence of the clone (rk1411) recovered above was determined as follows. By firstly using a kilo-sequence deletion kit (manufactured by TaKaRa, Co.), plural plasmid DNAs were prepared by subjecting the clone rk1411 to each deletion of about 300 bp from the single side thereof. The base sequences of the DNAs were determined by using an automatic sequencer (manufactured by Applied BioSystems). Additionally, a specific oligonucleotide primer for rk1411 was prepared; by using the automatic sequencer, the base sequences thereof were also analyzed from the opposite direction. Finally, the whole base sequence of rk1411 was determined. The base sequence is shown as SQ ID No. 3 in the sequence listing. Additionally, the amino acid sequence of the protein is shown as SQ ID No. 4.

Example 2 (Identification of the function of rk1411)

[0046]

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(1) By using T7 RNA polymerase, cRNA (RNA complementary to cDNA) was prepared in vitro from the plasmid carrying the clone (rk1411) as described above (see Sekine, T., et al. J. Biol. Chem., Vol. 272, pp. 18526-9, 1997).

According to the method already reported (Sekine, T., et al. J. Biol. Chem., Vol. 272, pp. 18526-9, 1997), the resulting cRNA was injected in the <u>Xenopus</u> oocyte; the oocyte was subjected to an uptake test with various radiolabeled organic anions and organic cations. As shown in Fig. 1, consequently, the oocyte in which rk1411 was expressed could take up ¹⁴C-PAH (p-aminohippuric acid), ³H-ochratoxin A and ³H-estrone sulfate. Alternatively, the oocyte never transported one typical organic cation ¹⁴C-TEA (tetraethylammonium).

The organic anion transport with rk1411 was subjected to the Michaelis-Menten dynamic test. By examining the change in the uptake of PAH, estrone sulfate and ochratoxin A at various concentrations, the dependency of the rk1411 transport on the concentrations of these substrates was examined. The uptake experiments of radiolabeled PAH, estrone sulfate and ochratoxin A were carried out by using the occyte injected with rk1411 cRNA according to the method described above. The results are as follows (see Fig. 2): the Km values of PAH, estrone sulfate and ochratoxin A were $4.7~\mu$ M, $2.3~\mu$ M and $0.74~\mu$ M, respectively. The results are shown below in Table 1.

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Table 1

Results of Michaelis-Menten dynamic test										
	Km (μM)	Vmax (pmol/hr/oocyte)	Vmax/Km (μl/hr/oocyte)							
PAH	64.7 ±10.0	23.3 ± 2.8	0.360							
Estrone sulfate	2.34 ± 0.20	7.60 ± 0.44	3.24							
Ochratoxin A	0.739 ± 0.178	3.08 ± 0.33	4.17							

(2) So as to examine the substrate selectivity of rk1411, various anionic substances were added to the ³H-estrone sulfate uptake experiment system with the occyte injected with rk1411 cRNA, to examine their influences (inhibition experiment). The ³H-estrone sulfate uptake experiment was conducted by using the occyte injected with rk1411 cRNA according to the method described above. In the presence and absence of 1 mM each compound (with no label), the uptake of ³H-estrone sulfate was assayed. Consequently, various anionic substances (taurocholic acid, cholic acid, bromosulfophthalein, probenecid, indocyanine green, burnetanide, cefoperazone, pyroxicam, furosemide, azidothymidine, benzylpenicillin and the like) significantly inhibited the ³H-estrone sulfate transport with rk1411 (see Fig. 3). Meanwhile, cationic substances such as tetraethylammonium, guanidine, quinidine and berapamil never exerted any such inhibitory action (see Fig. 3). The results indicate that rk1411 is a multi-selective transporter and primarily recognizes organic anions. Hence, rk1411 was designated OAT3 (organic anionic transporter 3) as a third member of the OAT family.

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Example 3

[0047] The expression of the OAT3 gene in individual rat tissues was analyzed (Northern blotting). The OAT3 cDNA in the whole length was labeled with ³²P-dCTP; by using the resulting cDNA as probe, RNAs extracted from various rat tissues were subjected to Northern blotting as follows. 3 µg of poly(A) + RNA was electrophoresed on 1 % agarose/formaldehyde gel and subsequently transferred on a nitrocellulose filter. The filter was hybridized overnight in a hybridization solution containing the whole length of the ³²P-dCTP-labeled OAT3 cDNA at 42 °C. The filter was rinsed in

0.1 x SSC containing 0.1 % SDS at 65 °C.

[0048] The Northern blotting results (see Fig. 4) indicate that a strong band was detected around 2.4 Kb in the RNAs from the kidney, liver and brain. Visually weak expression was also observed in the eyes.

5 Example 4

[0049] Because OAT3 was most strongly expressed in brain among the members of the OAT family, an attempt was made to deduce the role thereof in brain at an inhibition experiment of the OAT3 transport with various metabolites of neurotransmitters (mainly organic anions). As shown in Fig. 5, noradrenalin and serotonin metabolites inhibited the OAT3 transport of estrone sulfate, suggesting a possibility that these metabolites per se might be substrates of OAT3. The evidence indicates that OAT3 has an action to excrete neurotransmitter metabolites out of brain as one function of cerebral type OAT3.

Example 5

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Isolation and analysis of human-type multi-selective organic anion transporter 3 (OAT3) cDNA

[0050] EST (expressed sequence tag) data base was screened by using the rat OAT3 cDNA isolated previously by the inventors. Human EST clone (H20345) with high homology to the rat OAT3 was identified. A part (333 bp) of the base sequence of the clone was synthesized by PCR. The cDNA fragment was labeled with ³²P, which was then used as probe for the following screening.

[0051] The human kidney cDNA library maintained by the inventors was subjected to screening with the probe. Hybridization was effected all day long and overnight in a hybridization solution at 37 °C; subsequently, the filter membrane was rinsed in $0.1 \times SSC/0.1$ % SDS at 37 °C. As the hybridization solution, use was made of a buffer, pH 6.5, containing 50 % formamide, $5 \times SSC$ (standard saline citrate), 3 x Denhard solution, 0.2 % SDS, 10 % dextran sulfate, 0.2 mg/ml modified salmon sperm DNA, 2.5 mM sodium pyrophosphate, 25 mM MES, and 0.01 % Antifoam B (manufactured by Sigma, Co.). The clone isolated in λ ZipLox was further subcloned in a plasmid vector pZL by in vitro excision method. Consequently, a novel human organic anion transporter 3 (hOAT3) with an organic anion transport activity was recovered. The analysis of the transport function thereof is described below in Example 6.

[0052] The base sequence of hOAT3 was determined by the following method. Oligonucleotide primers specific to hOAT3 were sequentially synthesized. By using an automatic sequencer (manufactured by Applied BioSystems, Co.), the base sequence was analyzed, starting from both the 5'- and 3'-termini. Finally, the whole base sequence of hOAT3 was determined. The determined base sequence is shown as SQ ID No. 1 in the sequence listing. Based on the cDNA sequence, the amino acid sequence encoding hOAT3 is described as SQ ID No. 2 in the sequence listing.

[0053] The base sequence of the cDNA is shown in Table 2, while the amino acid sequence is shown in Table 3, in a corresponding manner.

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Table 2

Base sequence of hOAT3 cDNA

CTEAGCTGCC CTACTACAGC AGCTGCCGGC CCCTAGGACA GAGCAGGGAC CTCAACTACA CTEATCACCA ECCCCATCGE ATCCAGACCC EGCCACCAGC TCTEGCTCGT CTTECCCCAG TECCATGACC TTCTCGGAGA TCCTGGACCS TGTGGGAAGC ATGGGCCATT TCCAGTTCCT SCATGTASCC ATACTEGGCC TCCCGATCCT CAACATGGCC AACCACAACC TGCTGCAGAT CTTCACAGCC GCCACCCCTG TCCACCACTG TCGCCCGCCC CACAATGCCT CCACAGGCC TTEGGTECTC CCCATGGGCC CAAATGGGAA GCCTGAGAGG TGCCTCCGTT TTGTACATCC SCCCAATECC AGCCTEGGCA ATEACACCCA GAGGGCCATE GAGCCATECC TGGATGGCTG GETCTACAAC AGCACCAAGE ACTCCATTET GACAGAGTEG GACTTEGTET GCAACTCCAA CANACTGAAG GAGATGGCCC AGTCTATCTT CATGGCAGGT ATACTGATTG GAGGGCTCGT ECTTEGAGAC CTETCTGACA SETTTGGCCG CAGGCCCATC CTGACCTGCA GCTACCTGCT GCTGGCAGCC AGCGGCTCCG GTGCAGCCTT CAGCCCCACC TTCGCCATCT ACATGGTCTT CCGCTTCCTG TGTGGCTTTG GCATCTCAGG CATTACCCTG AGCACCGTCA TCTTGAATGT EGAATEGETE CCTACCCEEA TECGGECCAT CATETCEACA GCACTCEEGT ACTECTACAC B 1 0 CTTTEGCCAG TTCATTCTEC CCSECCTEGC CTACGCCATC CCCCAGTGGC GTTEGCTGCA STTAACTETE TOCATTOCCT TOTTOETOTT CTTCCTATCA TOCTEETEGA CACCAGAGTO CATACGCTEG TTEGTCTTET CTEGAAGTC CTCEGAGECC CTGAAGATAC TCCGGCGGGT SECTETCTTC AATGGCAAGA AGGAAGAGGS AGAAAGGCTC AGCTTEGAGG AGCTCAAACT CAACCTGCAG AAGGAGATCT CCTTGGCCAA GGCCAAGTAC ACCGCAAGTG ACCTGTTCCG GATACCCATG CTGCGCCGCA TGACCTTCTG TCTTTCCCTG GCCTGGTTTG CTACCGGTTT

	1.150	1160	1170	1180	1190	1200
		AGTTTGGCTA				•
5	1210	1220	1230	1240	1250	1260
		GETGGGGTCG				
	1270	1280	1290	1300	1310	1320
		CATACCACTC				
10	1330	1340	1350	1360	1370	1380
	TCTCACCTTT	GTGCCCTTGG	ACTIGCAGAC	CGIGAGGACA	GIALIGGUIG	AADDIIIDI
	1390	1460	1410	1420	1430	1440
	EGGATGCCTA	TCCAGCTCCT	TCAGCTGCCT	CTTCCTCTAC	ACAAGTGAAT	TATACCCCAC
15	1450	·· 1460	1470	1480	1490	1500
	AETCATCAGG	CAAACAGGTA	TGGGCGTAAG	TAACCTGTGG	ACCCGCGTGG	GAAGCATGGT
	1510	1520	1530	1540	1550	1560
		STGAAAATCA	•			
20	1570	1580	1590	1600	16-10	1620
	GATCACCGCC	CTCCTCGGGG	6CAGT6CT6C	CCTCTTCCTS	CCTGAGACCC	TGAATCAGCC
	1630	1640	1650	1660	1670	1680
25		•				AGCCAAAGCA
	1690	1700	1710	1720	1730	1740
		-				GACCAGGCCT
	1750	1760	1770	1780	1790	1800
30						GATCCTAGCC
	1810	1820	1830	1840	1850	1860
						TGAAGCCTTC
	187	1880	1890	1900	1910	1920
35						AGCCCTGGCC
	193	1940	1950	1860	1976	1980
			•			CTGCCATTCT
	199	0 2000	2010	2020	2030	2040
40						TTCCCCTGAG
	205	0 2080	207	8 208i	2090	2100
			-			S ATGAGAAGTC
45	211	0 212	213	0 214	D 215	2150
						: AGCGATAAGA
	217	0 218	0 219	0 220	0 221	0 2220
	ACTUTARAA	KAAAAAAA.				

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Table 3

Base sequence of hOAT3 amino acid

	5'	ATG ACC	133 TTC TC6	A DAG	42 TC CT6	GAC	151 CST	6T6		160 AGC	ATG		169 CAT	स्तर	CAG	178 TTC
10		Ret Thr	Phe Ser	Glu I	le Leu	Åsp	Arg	Val	Gly	Ser	Me t	Gly	His	Phe	6 l n	Pho
		CTG CAT	187 GTA GCC	ATA C	96 TG GGC	стс	205 CCG	ATC		214 AAC	ATG		223 AAC	CAC	AAC	232 CTG
15		Lou His	Val Ala	lle L	en ela	Leu	Pro) [e	Lou	Asn	Met	E J A	Ásn	His	Asn	Leu
		CTG CAG	241 ATC TTC		50 CC 6CC	AGC	259 CCT	GTC	CAC	268 CAC	TGT	CGC	271 CCG	ccc		286 AAT
20		Lau Gir	lle Phe	The A	la Ala	Thr	Pro	Val	Hi s	His	Cys	Arg	Pro	Pro	HIS	Asn
		GCC TCC	295 ACA 666		GG GTS	CTC	313 CCC	AT6		322 CCA	AAT	333	331 AAG	CCT	GAG	340 AGG
		Ala Sei	The 61	Pro T	rp Val	Leu	Pro	Me t	EIA	Pro	Asn	GIY	LYE	Pra	614	a ı A
25		TGC CT	349 CGT TT		158 AT CCG	ccc	367 AAT	GCC	AGC	376 CTG	ccc	AAT	385 GAC	ACC	CAĠ	384 AGG
		Cys La	Arg Ph	Val H	ijs Pro	Pro	Asn	Ala	Ser	Leu	Pro	Asn	Asp	Thr	6 I n	Arg
30		GCC AT	403 GAG CC		112 CTG GAT	GGC	421 TGG	GTC	TAC	0E4	AGC	ACC	439 AA6	GAC	TCC	448 ATT
		Ala Me	t Glu Pr	Cys L	Lau Asp	Gly	Trp	Vel	Tyr	Υял	Ser	Thr	Lys	Asp	Ser	Ha
35		GTG AC	457 A GAG TG		166 FTG GTG	TEC	AAC		AAC	484 AAA	ств	AAG	493 6A6		ecc	502 CAG
		Yai Th	r Glu Tr	p Asp I	Lou Val	Çys	Asn	. Ser	Asn	LYS	Lea	Lys	6 i u	Met	Aja	Gin
40		TCT AT	C TTC AT			CTG	529 ATT		666	53B CTC		CTT	54T GGA			556 TCT
40		Ser 11	e Phe Me	t Ala (61y 11e	Leu	110	. G1 A	Gly	Leu	Val	Leu	6 3 Y	Asp	Lou	Ser
		GAC AG	565 6 TTT 66	c cec	574 AGG CC		583 CT6	ACC	TEC	592 AGC		CTG	601 CTG		GCA	610 6CC
45		Asp Ar	g Pha Gi	y Årg 4	Arg Pro	lle	Leu	The	CAR	Şer	TYE	Fea	Leu	Leu	. Ala	g) Å
		YEC EE	519 C TCC 66	T GCA	628 GCC TTC			ACC			ATG			ETC		
50		Ser Gl	y Ser 61													
		TTC CT	673 6 T6T 66		682 660 AT	TCA	691 660		ACC	CTE		ACC	709 GTC		TTG	718 TAA
		Phe Le	CYS SI	y Phe	Sly II	Ser	Gly	H	Thr	Leu	Se r	Thr	Vel	110	Lau	nek

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	GTG	EAA	727 TGG	GTG	CCT	736 ACC	CGG	ATE	CGG	GCC	ATC		TCG	ACA	GCA	CTC	GG6	772 TAC
5	leV	61 u	Trp	Val	Pro	Thr	Arg					Met				Leu	Gly	Tyr
	TEC	TAC	781 ACC	TTT	GGC	790 CAG	TTC	ATT	799 CTG	ccc	GGC	808 CTG	600	TAC	8 1 7 6CC	ATC	ccċ	826 CAG
10	Cys	 Туг	The	Phe	GIY	Gin	Phie	lie	Leu	Pro	GIY	Leu	Ala	Tyr	Ala	lle	Pro	Gin
		·	835			844			853			862			871			880
	TEG	732 	TGG	CTG			ACT					277	TTC	GTC	TTC	TTC	CTA	TCA
15	Trp	Arg	Trp	Lau	a 19	Lau	Thr	Val	Ser	110	Pro	Phe	Phe	Vai	Pha	Phe	Leu	Ser
	TCC	TGG	889 TGG				TCC						TTG			YAG		934 TCG
	Ser	Trp	Trp	Thr	Pro	Glu	Ser	lle	Arg	Trp	Lau	Va l	Leu	Ser	Ely	LYS	Ser	Ser
20	GAG	GCC	943 CTG			CTC	CEG	CEE	ETG	GCT	GTC	TTC	AAT			AAG		988 6AG
	Glu	Ala	Lau	Lys			Arg						Asa	Gly	Lys	Lys	Glu	Glu
25 .	GGA		997 AGG		AGC	1006 TTG	GAG	6AG	1015 CTC	AAA	стс	1024 AAC	ств	CAG	EEG 1 DAA	GAG	ATC	1042 TCC
	GIY	611	Arg	Leu	Ser	Leu	610	61 u	Leu	Lys	Løu	Asn	Leu	Gin	Lys	Glu	110	Ser
30	TTG		1 05 1 AAG		AAG	1 0 6 0 TAC	ACC	GCA	1069 AGT	SAC	CTG	1078 TTC	CES		1087 CCC			1096
	Lau	SIA	Lys	Ala	Lys	Tyr	Thr	Ala	Ser	Asp	Leu	Phe	Arg	ite	Pro	Met	Leu	Arg
35	cec	ATG	1105 ACC		TGT	1114 CTT	TCC	CTG	1123 600	TGG	TTT	1132 GCT	ACC		1141 TTT			1150 TAT
	D 1 A	Het	Thr	Phe	Cys	Leu	Ser	Leu	Ala	Trp	Phe	Ala	Thr	GIY	Pha	Ala	Tyr	Tyr
			1159			1168	~	C 4 8	1177	CCA		1186			1195	CTE		1284
40							GAA				~							
	561	. Fea			-							1240						110
45	ATC	ידד:	1213	686	670	GAT	GTC	CCA	S GCC	AAG	TT	ATC	ACC	ATO	CTC	TCC	TTA	
	116	Phe	SI	r ely	Yal	Asp	Yal	Pre	Ala	Lys	Ph	: !16	Th	114	Lac	Sar	Lau	Ser
	710		1267						1285			1294			1303		. ee	1312 : GCC
50										-								Als
	. , ,			, ,,,,,,,	1112	,,, ,		011								;	, 017	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,

		1	321		1	330		•	1339		1	348		1	357		1	366
		TTG	GCT	CTC	ACC	TTT	GTS	CCC	TIG	gac	TTG	CAG	ACC	6T6	AGG	ACA	GTA	TTG
5											Lau							
			375		1	384		:	1393			402		1	411		1	420
	GCT	GTG	TTT	666	AAG	GGA	TGC	CTA	TCC	Yec	TCC	TTC	Yec	TGC	CTC	TTC	CTC	TAC
10											Ser							Tyr
			429		1	438			1447		1	456		1	465		1	474
	ACA	AGT	eyy	TTA	TAC	CCC	YCY	STC	ATC	YEE	CAA	YCY	T39	ATG	eec	GLY	AGT.	AAC
15	Thr	Ser	8 i v	Lau	Туг	Pro.	Thr	Vai	He	Arg	6 i n	Thr	Giy	Met	6 l y	Va 1	Ser	Asn
			1483		1	492			1501		1	510		1	1519		1	528
	CTE	TGG	YCC								222							
	Leu	Trp	Thr								Pro							
20		!	1537		1	546			1555		1	564		•	1573		1	582
											666			GCC				
											GIY			Ala		Leu		
25			1201						1600			1212			1577			636
		GCT	ecc	CTC		CTG	CCT	6AG	ACC	CTG	AAT	CAG	ccc	TTG	CCA		ACT	
											Åsn							11.
30		GAC	CTE	GAA	AAC	T6G	TCC	CTG	ÇEG	GCA	AAG							
					Asn						Lys	Lys	Pro	Lys	6) n	Giu	Pre	Giu
35											CAG							
																		61 y
40		AGC		3'														
	Ser	Şer	**															

Example 6

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Identification of hOAT3 function

[0054] By using T7 RNA polymerase, cRNA (RNA complementary to cDNA) was prepared in vitro from the plasmid comprising the hOAT3 recovered above by the method by Sekine, et al. (see Sekine, T., et al. J. Biol. Chem., Vol. 272, pp. 18526-9, 1997).

[0055] According to the already reported method of Sekine, et al. (Sekine, T., et al. J. Biol. Chem., Vol. 272, pp. 18526-9, 1997), the resulting hOAT3 cRNA was injected in the Xenopus oocyte; the oocyte was subjected to an uptake test with various radiolabeled organic anions and organic cations. The control oocyte cell (oocyte cell with no injection of hOAT3 cRNA) and the oocyte cell injected with hOAT3 cRNA were cultured in a buffer containing the following radiolabels for one hour, to assay the uptake of the radiolabels into the oocytes.

[0056] The results are shown in Figs. 6 to 18. In each figure, open column expresses the case of the control oocyte used; and closed column expresses the case of the oocyte injected with hOAT3 cRNA. Fig. 6 depicts the uptake activity of ¹⁴C-PAH (p-aminohippuric acid) (10 µM); Fig. 7 depicts the uptake activity of ³H- strone sulfate (50 nM); Fig. 8

depicts the uptake activity of 3 H-dehydroepiandrosterone sulfate (50 nM); Fig. 9 depicts the uptake activity of 3 H-ochratoxin A (100 nM); Fig. 10 depicts the uptake activity of 3 H-cimetidine (150 nM); Fig. 11 depicts the uptake activity of 3 H-estradiol glucuronide (50 nM); Fig. 12 depicts the uptake activity of 3 H-prostaglandin E2 (1 nM); Fig. 13 depicts the uptake activity of 14 C-faurocholic acid (10 μ M); Fig. 14 depicts the uptake activity of 14 C-glutaric acid (10 μ M); Fig. 15 depicts the uptake activity of 3 H-methotrexate (100 nM); Fig. 16 depicts the uptake activity of 14 C-salicylic acid (1 μ M); Fig. 17 depicts the uptake activity of 14 C-indomethacin (10 μ M); and Fig. 18 depicts the uptake activity of 14 C-cholic acid (10 μ M).

[0057] As shown in these figures, the values of these radiolabels in the oocyte with hOAT3 expression were higher than the values thereof in the control oocyte, suggesting that hOAT3 transported these compounds.

[0058] Consequently, the oocyte with hOAT3 expression takes up ¹⁴C-PAH (p-aminohippuric acid), ³H-estrone sulfate, ³H-dehydroepiandrosterone sulfate, ³H-ochratoxin A, ³H-cimetidine, ³H-estradiol glucuronide, ³H-prostaglandin E2, 14C-taurocholic acid, ¹⁴C-glutaric acid, ³H-methotrexate, ¹⁴C-salicylic acid, ¹⁴C-indomethacin, and ¹⁴C-cholic acid. On contrast, hOAT3 never transported the typical organic cation ¹⁴C-TEA (tetraethylammonium) (not shown in the figures).

[0059] Then, the hOAT3 transport of organic anions was examined at the Michaelis-Menten kinetic test. By examining the change in the hOAT3 uptake of estrone sulfate and methotrexate at various concentrations, the dependency of the OAT3 transport on the concentrations of these substances was examined. The uptake experiment of radiolabeled estrone sulfate and methotrexate was carried out by using the oocyte injected with hOAT3 cRNA and the control oocyte (with no injection of cRNA), by the method described above. Consequently, the Km values of estrone sulfate and methotrexate were 3.08 μM and 2.22 μM, respectively.

[0060] So as to examine the substrate selectivity of hOAT3, various anionic substances were added to the ³H-estrone sulfate uptake experiment system with the oocyte injected with hOAT3 cRNA, to examine their influences (inhibition experiment).

[0061] The ³H-estrone sulfate uptake experiment was conducted by using the oocyte injected with hOAT3 cRNA according to the method described above.

[0062] More specifically, the control oocyte (oocyte with no injection of hOAT3 cRNA) and the oocyte with injection of hOAT3 cRNA were cultured in a buffer containing 50 nM ³H-estrone sulfate alone or containing non-radiolabeled compounds at 500 µM or the concentration shown in the figure for one hour, to assay the uptake of ³H-estrone sulfate. When the uptake of 50 nM ³H-estrone sulfate singly contained in the buffer into the oocyte with injection of hOAT3 cRNA was designated 100 %, the individual uptake values in the buffer containing inhibitory agents were expressed in the buffer

[0063] The results are shown in Fig. 19. As shown in Fig. 19, all these compounds inhibited the uptake of ³H-estrone sulfate into the oocyte injected with hOAT3 cRNA, indicating that these compounds were interactive with hOAT3. Consequently, it was indicated that various anionic substances (estrone sulfate, PAH, taurocholic acid, probenecid, furosemide, zidovudine, penicillin G, BSP, glutaric acid, indomethacin, and methotrexate) significantly inhibited the transport of ³H-estrone sulfate with hOAT3 (see Fig. 19). Alternatively, tetraethylammonium as one of typical organic cations never exerted any inhibitory action. Based on these results, it is evidenced that the inventive hOAT3 is a multi-selective organic anion transporter.

40 Industrial Applicability

[0064] The invention provides a novel organic anion transporter with wide substrate selectivity of organic anions and in selective distribution in brain and liver and the like.

[0065] The inventive organic anion transporter is involved in the uptake of various drugs in cells and is also involved in the dynamics of drugs in biological organisms. Therefore, the inventive organic anion transporter is useful not only for the cell viability and activation but also for the screening of pharmacokinetics.

Claims

- 1. A cerebral type organic anion transporter OAT3.
- 2. A cerebral type organic anion transporter OAT3 according to claim 1, wherein the cerebral type organic anion transporter OAT3 is of an amino acid sequence represented by SQ ID No. 2 or 4 in the sequence listing or of an amino acid sequence with such a modification of the amino acid sequence represented by SQ ID No. 2 or 4 as deletion of a part of the amino acid sequence, or substitution or addition with other amino acids.
- 3. A nucleic acid encoding a protein of an amino acid sequence represented by SQ ID No. 2 or 4 in the sequence

listing or of an amino acid sequence with such a modification of the amino acid sequence represented by SQ ID No. 2 or 4 as del tion of a part of the amino acid sequence, or substitution or addition with other amino acids.

- A nucleic acid according to claim 3, wherein the nucleic acid is DNA of a base sequence repr sented by SQ ID No. 1 or 3 in the sequence listing.
 - 5. A nucleic acid comprising at least 14 consecutive nucleotides in series of the DNA of a base sequence represented by SQ ID No. 1 or 3 or a strand complementary to the 14 consecutive nucleotides in series.
- 6. A nucleic acid according to claim 5, wherein the number of the nucleotides is 20 or more.
 - 7. An antibody capable of recognizing a cerebral type organic anion transporter OAT3 according to claim 1 or 2.

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Figure 1

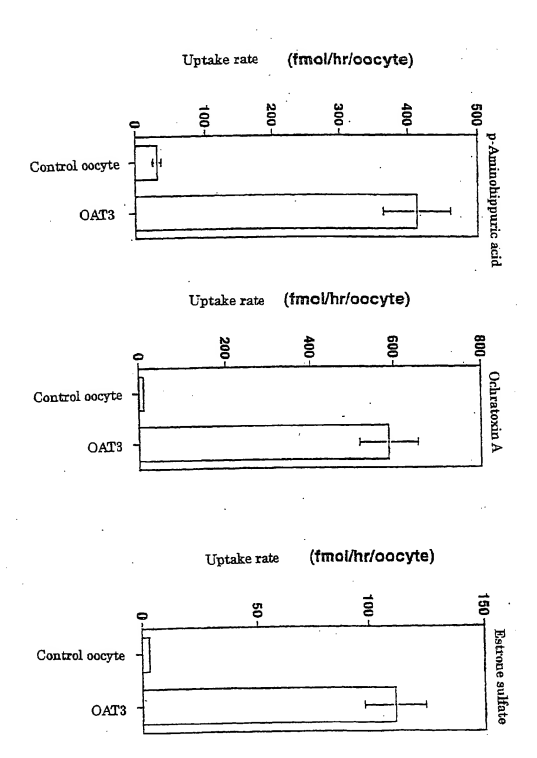


Figure 2

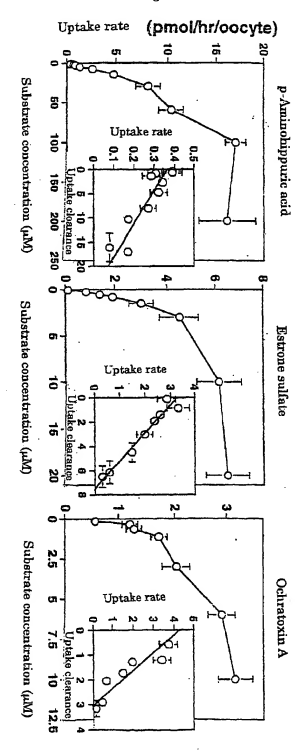
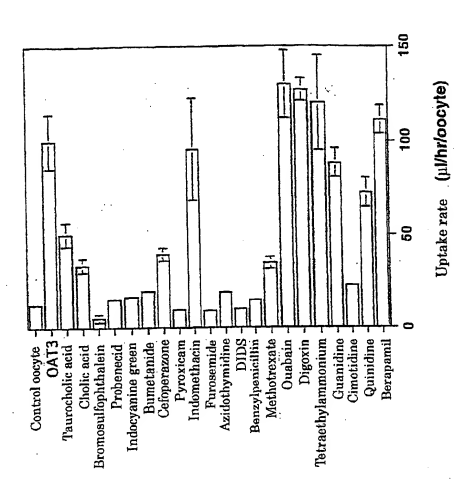


Figure 3



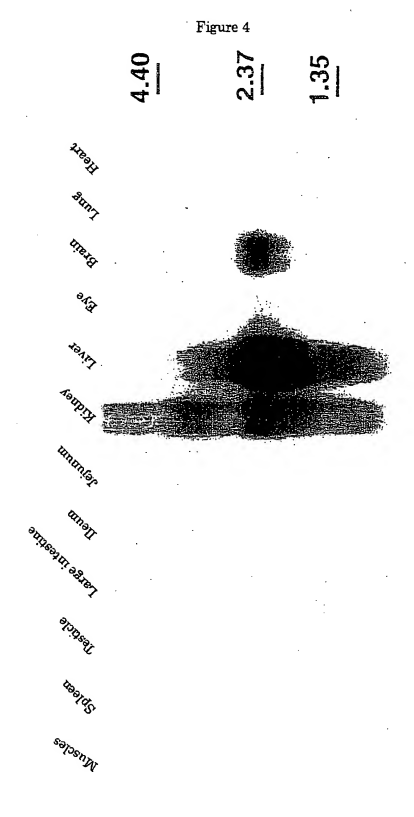


Figure 5

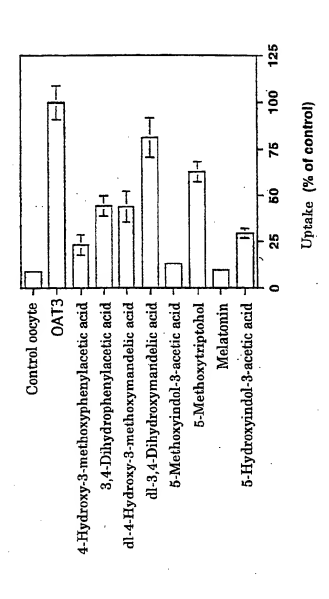


Figure 6

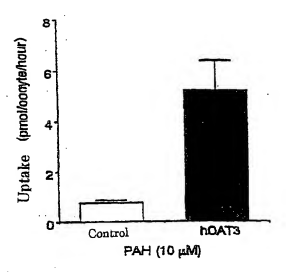


Figure 7

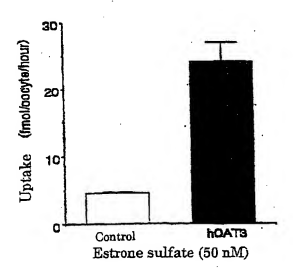
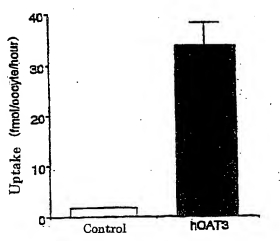


Figure 8



Dehydroepiandrosterone sulfate (50 nM)

Figure 9

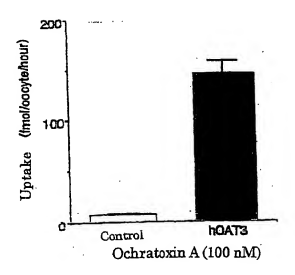


Figure 10

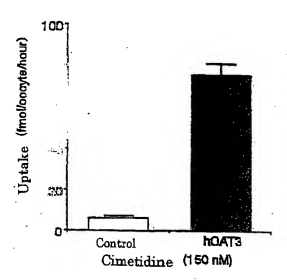


Figure 11

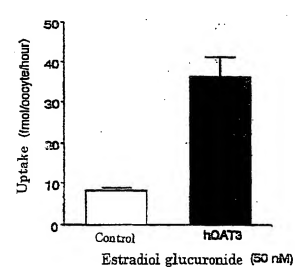


Figure 12

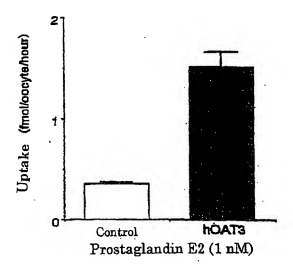


Figure 13

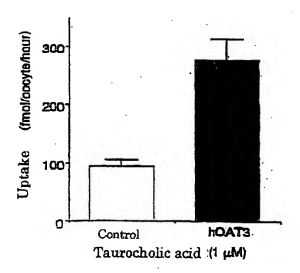


Figure 14

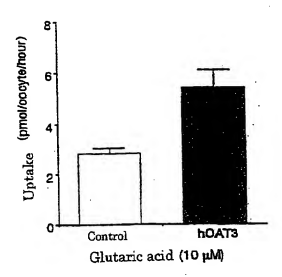


Figure 15

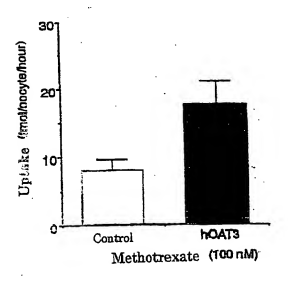


Figure 16

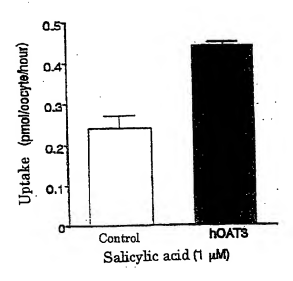


Figure 17

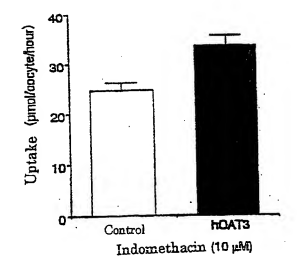


Figure 18

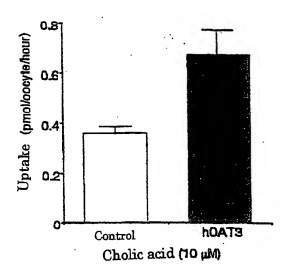
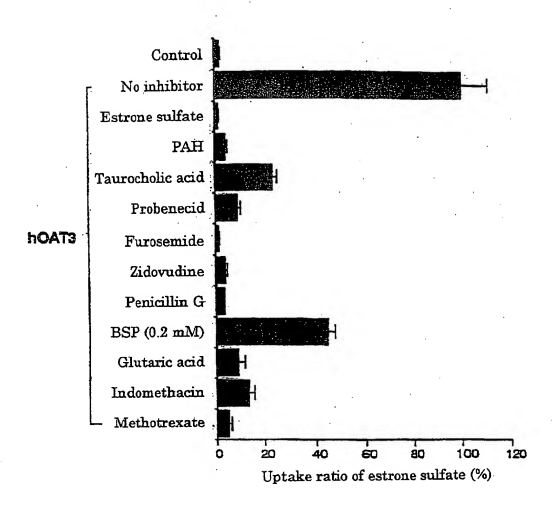


Figure 19



INTERNATIONAL SEARCH REPORT International application No. PCT/JP99/05120 -

A. CLASSIFICATION OF SUBJECT MATTER Int.Cl ⁷ C07K14/435, C07K16/18, C12	2P21/02, C12P21/08, C12N1	15/12										
According to International Patent Classification (IPC) or to both national classification and IPC												
B. FIELDS SEARCHED												
Minimum documentation searched (classification system followed by classification symbols) Int.Cl ⁷ C07K14/435, C07K16/18, C12P21/02, C12P21/08, C12N15/12												
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched												
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) WPI, WPI/L, BIOSIS PREVIEW, CAS ONLINE, GenBank/EMBL/DDBJ/Geneseq												
C. DOCUMENTS CONSIDERED TO BE RELEVANT												
Category* Citation of document, with indication, where ap		Relevant to claim No.										
of a New Multispecific Organic Ar	Endou H. et.al., "Molecular Cloning and Characterization of a New Multispecific Organic Anion Transporter from Rat Brain", J. Biol. Chem. (May, 1999), Vol. 274, No. 19,											
anion transporter 2 expressed pre	Endou H. et. al., "Identification of multispecific organic anion transporter 2 expressed predominantly in the liver", FEBS Letters (June, 1998), Vol. 429, pages 179-182											
Further documents are listed in the continuation of Box C.	See patent family annex.											
Special categories of cited documents: A** document defining the general state of the art which is not considered to be of particular relevance earlier document but published on or after the international filing date "L** document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "O** document referring to an oral disclosure, use, exhibition or other means "P** document published prior to the international filing date but later than the priority date claimed Date of the actual completion of the international search	"I" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art document member of the same patent family											
21 December, 1999 (21.12.99) Name and mailing address of the ISA/	Date of mailing of the international search report 28 December, 1999 (28.12.99) Authorized officer											
Japanese Patent Office	Telephone No.											

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